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A1 20030911 (200367)

WO 2003057924 A1 WO 2003-US180 20030103; US 2003171325 A1 Provisional US

AU 2003207448 A1 20030724 (200421)

US 2003171325

ADT

2002-345099P 20020104, US 2003-336610 20030103; AU 2003207448 A1 AU 2003-207448 20030103

FDT AU 2003207448 A1 Based on WO 2003057924

PRAI US 2002-345099P 20020104; US 2003-336610 20030103

AN 2003-697277 [66] WPIDS

AB W02003057924 A UPAB: 20031014

NOVELTY - Solid-phase oligonucleotide synthesis (M1) comprising capping the ends of a double stranded DNA (dsDNA) consisting of sense and antisense oligonucleotides, cleaving the dsDNA at or near a Watson-Crick base pair mismatch and digesting the uncapped dsDNA, is new.

DETAILED DESCRIPTION - Solid-phase oligonucleotide synthesis (M1) comprising:

(a) synthesizing a sense and an antisense oligonucleotide;

- (b) annealing the sense and antisense oligonucleotides to form double stranded DNA (dsDNA);
 - (c) capping the ends of the dsDNA;
- (d) cleaving the dsDNA, where **cleavage** occurs at or near a Watson-Crick base pair mismatch; and
 - (e) digesting the uncapped dsDNA, is new. INDEPENDENT CLAIMS are also included for:
 - (1) forming long polynucleotides (M2) comprising:
- (a) synthesizing a first proofread dsDNA by employing the steps of $\mbox{M1};$
 - (b) synthesizing a second proofread dsDNA; and
- (c) ligating the first proofread DNA with the second proofread DNA to form a long polynucleotide;
 - (2) an apparatus for performing M1; and
 - (3) an apparatus for performing M2.

USE - M1 is useful for synthesizing high-fidelity oligonucleotides (claimed), which may be used as probes, or in medical diagnostics, life sciences, and the pharmaceutical industry. The sequences produced may be used as synthetic genes and synthetic chromosomes to direct protein synthesis in living systems, or for information storage in devices such as molecular computers.

ADVANTAGE - M1 is used to synthesize high purity short oligonucleotides with fewer errors and eliminates the need for inefficient HPLC or other cleanup.

Dwg.0/3

- L4 ANSWER 2 OF 17 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN
- AN 2003:567097 BIOSIS
- DN PREV200300568401
- TI **Solid-phase synthesis** of core 2 O-linked glycopeptide and its enzymatic sialylation.
- AU Takano, Yutaka; Kojima, Naoya; Nakahara, Yuko; Hojo, Hironobu; Nakahara, Yoshiaki [Reprint Author]
- CS Department of Applied Biochemistry, Institute of Glycotechnology, Tokai University, 1117 Kitakaname, Hiratsuka-shi, Kanagawa, 259-1292, Japan yonak@keyaki.cc.u-tokai.ac.jp
- SO Tetrahedron, (13 October 2003) Vol. 59, No. 42, pp. 8415-8427. print. ISSN: 0040-4020 (ISSN print).
- DT Article
- LA English
- ED Entered STN: 3 Dec 2003

Last Updated on STN: 3 Dec 2003

The core 2-type tetrasaccharide building blocks (1a/1b) for solid

-phase synthesis of glycopeptide were synthesized via

stereaselective glycosylation of the disaccharyl Ser/Thr (3a/3b) with a
glycosyl fluoride (2) carrying the 2-trichloroacetamido group that was
readily converted into a 2-acetamido group by reduction. A segment of
glycoprotein leukosialin (215-224) was synthesized by the solid-phase
protocol, the building block (1b) being utilized. Cleavage of
the synthetic glycopeptide from resin was effected with reagent K and
subsequent treatment of the product with a cocktail for the 'low-acidity

TfOH' facilitated complete removal of the benzyl groups with minimum loss of glycosidic linkages. To the deprotected glycopeptide (21), were enzymatically introduced N-acetylneuraminic acid (sialic acid) residues in remarkably high efficiency by using the specific sialyltransferases.

- L4 ANSWER 3 OF 17 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 2003:324010 CAPLUS
- DN 139:100670
- TI A Novel and Rapid Encoding Method Based on Mass Spectrometry for "One-Bead-One-Compound" Small Molecule Combinatorial Libraries
- AU Song, Aimin; Zhang, Jinhua; Lebrilla, Carlito B.; Lam, Kit S.
- CS Division of Hematology and Oncology Department of Internal Medicine, UC Davis Cancer Center, Sacramento, CA, 95817, USA
- SO Journal of the American Chemical Society (2003), 125(20), 6180-6188 CODEN: JACSAT; ISSN: 0002-7863
- PB American Chemical Society
- DT Journal
- LA English
- OS CASREACT 139:100670
- AB A method for the preparation and encoding of readily deconvoluted combinatorial libraries is discussed. Beads are prepared with topol. segregated regions an inner region to which is bound coding tags and an outer segment to which the library compound is bound. Coding blocks are attached to the inner resin by a cleavable methionine-containing linker; the coding blocks are chosen to have similar reactivities to the building blocks incorporated in the synthesis of the combinatorial library. Synthesis of the library leads to the functionalization of the library-containing portion of the resin bead and the coding portion of the resin bead.

Cleavage of the linkers for the coding blocks from the resin bead
 by Edman degradation with cyanogen bromide yields lactones whose mass is
determined

by FT-MALDI mass spectroscopy. Anal. of the lactones isolated from a given bead yields the mass of each of the fragments present; by careful choice of coding blocks and reactants, the identities of the building blocks incorporated into a library bead and of the library member attached to that bead can be readily derived from the fragment masses. A combinatorial library is prepared and tested for the binding of library members to streptavidin; seventeen of the compds. are found to bind strongly to streptavidin by a colorometric assay and identified unambiguously by the library encoding method described here.

RE.CNT 61 THERE ARE 61 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L4 ANSWER 4 OF 17 SCISEARCH COPYRIGHT 2004 THOMSON ISI on STN
- AN 2003:352461 SCISEARCH
- GA The Genuine Article (R) Number: 669ZK
- TI A versatile approach towards regioselective platinated DNA sequences
- AU Heetebrij R J; de Kort M; Meeuwenoord N J; den Dulk H; van der Marel G A; Van Boom J H (Reprint); Reedijk J
- CS Leiden Univ, Leiden Inst Chem, Gorlaeus Labs, Program 9502, NL-2300 RA Leiden, Netherlands (Reprint); Leiden Univ, Leiden Inst Chem, Gorlaeus Labs, NL-2300 RA Leiden, Netherlands
- CYA Netherlands
- SO CHEMISTRY-A EUROPEAN JOURNAL, (14 APR 2003) Vol. 9, No. 8, pp. 1823-1827. Publisher: WILEY-V C H VERLAG GMBH, PO BOX 10 11 61, D-69451 WEINHEIM, GERMANY.
 - ISSN: 0947-6539.
- DT Article; Journal
- LA English
- REC Reference Count: 30
 - *ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS*
- AB Undesired N-7 platination of 2'-deoxyguanosine residues at predetermined sites in an oligodeoxynucleotide (ODN) sequence is prevented by applying the sterically demanding diphenylcarbamoyl (DPC) as an

O-6-protecting group. The presence of a base-labile oxalyl linker between the immobilized 3'-nucleotide and controlled pore glass (CPG) allows cleavage of the protected ODN from the support and leaves DPC protection unaffected. This method provides an ODN with specifically blocked guanine-N-7 sites for platination. In the hexanucleotides prepared in this study, 5'-GGBGGT-3'(for B = T, C and A), a platinum GG adduct is introduced at G4,G5. These site-specific platinated hexamers were isolated in a yield of 65%, and were fully characterized by using reversed-phase HPLC (high performance liquid chromotography), LCMS (liquid chromatography-mass spectrometry), MALDI-TOF MS (matrix-assisted laser desorption/ionization time-of-flight mass spectrometry), PAGE (polyacrylamide gel electrophoresis) and Maxam-Gilbert sequencing analysis.

- L4 ANSWER 5 OF 17 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 2003:453923 CAPLUS
- DN 139:396154
- TI Towards the standard-module approach to disulfide-linked polypeptide nanostructures. I. Methodological prerequisites and mass spectrometric characterization of the test two-loop structure
- AU Mirgorodskaya, O. A.; Haselmann, K. F.; Kjeldsen, F.; Zubarev, R. A.; Roepstorff, P.
- CS Department of Chemistry, University of Southern Denmark, Odense, Den.
- SO European Journal of Mass Spectrometry (2003), 9(2), 139-148 CODEN: EJMSCL; ISSN: 1469-0667
- PB IM Publications
- DT Journal
- LA English
- Potentially biol.-active nanostructures can be created from single chains of unmodified peptides by crosslinking different regions of the chain by disulfide bonds and cleaving the chain at specified sites to obtain the final configuration. The availability of techniques for assembly and characterization of such structures was tested on a two-loop structure created from a 21-residue linear peptide. Directed intra-mol. disulfide bond formation was performed by inserting partial sequences favoring intra-mol. S-S bond formation ("loops") separated by partial sequences disfavoring such a process ("spacers") into the precursor sequence. Peptide bond cleavage by partial acid hydrolysis at specific sites (GG, NP/DP) inside the loops opened them; the same process in the spacer separated the loops. Synthesis, oxidation and bond cleavage were monitored by matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI ToF MS). The hydrolysis fragments of the produced nanostructures were characterized by tandem electrospray ionization Fourier transform mass spectrometry (ESI FT-MS) with collisional and electron capture dissocns. The latter technique was especially useful as it cleaves S-S bonds preferentially. feasibility of the proposed synthesis approach and the adequacy of the anal. techniques for the test structure were demonstrated.
- RE.CNT 19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L4 ANSWER 6 OF 17 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 2002:768751 CAPLUS
- DN 138:4807
- TI A New Method for the Preparation of Unprotected Peptides on Biocompatible Resins with Application in Combinatorial Chemistry
- AU Pastor, Jose J.; Fernandez, Irene; Rabanal, Francesc; Giralt, Ernest
- CS Institut de Recerca Biomedica de Barcelona Parc Cientific de Barcelona, Universitat de Barcelona, Barcelona, 08028, Spain
- SO Organic Letters (2002), 4(22), 3831-3833 CODEN: ORLEF7; ISSN: 1523-7060
- PB American Chemical Society
- DT Journal
- LA English

AB A synthetic strategy for the preparation of side chain free (deprotected) peptides on biocompatible solid supports is described. Final peptide detachment is afforded in mild basic conditions with no presence of scavengers or other additives, thus allowing single peptide-resin beads to be cleaved in mass spectrometry sample plates for direct sequencing using MALDI-TOF post-source decay. Using a [9-(hydroxymethyl)-9H-fluoren-2-yl]-carbamic acid moiety (I) as bifunctional linker, cleavable under mild basic conditions (morpholine in DMF), the final cleavage can be accomplished directly on MALDI plates to ensure no losses of pos. compds. This methodol. offers clear advantages for the development of one-bead-one-compound combinatorial libraries in addition to parallel and regular synthesis of peptides.

RE.CNT 18 THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 7 OF 17 MEDLINE on STN

AN 2002680717 MEDLINE

DN PubMed ID: 12440863

TI Oligonucleotides incorporating 7-(aminoalkynyl)-7-deaza-2'deoxyguanosines: duplex stability and phosphodiester hydrolysis by exonucleases.

AU Rosemeyer Helmut; Ramzaeva Natalya; Becker Eva-Maria; Feiling Elisabeth; Seela Frank

CS Laboratorium fur Organische und Bioorganische Chemie, Institut fur Chemie, Fachbereich Biologie/Chemie, Universitat Osnabruck, Barbarastrasse 7, D-49069 Osnabruck, Germany.

SO Bioconjugate chemistry, (2002 Nov-Dec) 13 (6) 1274-85. Journal code: 9010319. ISSN: 1043-1802.

Ι

CY United States

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 200305

ED Entered STN: 20021121

Last Updated on STN: 20030517

Entered Medline: 20030516

Oligonucleotides containing 7-(omega-aminoalkyn-1-yl)-7-deaza-2'-deoxyguanosines (1a-c) were investigated regarding their thermal stability (T(m) values) as well as their phosphodiester hydrolysis catalyzed by exonucleases. Those derivatives are suitable for the labeling of nucleic acid constituents as well as for the postlabeling of DNA. For this, the phosphoramidites 7a,c (obtained from the nucleoside 1a,b), protected by an isobutyryl group at the 2-amino group and a phthaloyl residue at the side-chain amino function, were synthesized. Using compounds 7a,c together with the phosphoramidite of 1c in solid-phase synthesis, a series of self-complementary and non-self-complementary oligonucleotides were prepared and characterized by MALDI-TOF mass spectrometry. A comparison of the T(m) values of the modified oligomers shows that the thermal stability of the duplexes

decreases with the length of the nucleobase 7-(omega-aminoalkyn-1-yl) side chain. Exonucleolytic cleavage of oligonucleotide single strands incorporating either the 7-(3-aminopropyn-1-yl) - or the 7-(4-aminobutyn-1-yl)-substituted nucleosides 1a or 1b, respectively, reveals that 3' --> 5' specific snake venom phosphodiesterase liberates 1a 5'-monophosphate but not the methylene-extended 1b 5'-monophosphate. On the contrary, the 5' --> 3' specific bovine spleen exonuclease is able to cleave off single 1a and 1b 3'-monophosphate residues; its action is, however, terminated in the case of oligonucleotides containing two consecutive 1a or 1b nucleotide units.

- L4 ANSWER 8 OF 17 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 2002:231045 CAPLUS
- DN 137:33511
- TI High-throughput peptide synthesis and peptide purification strategy at the low micromol-scale using the 96-well format
- AU Pipkorn, R.; Boenke, C.; Gehrke, M.; Hoffmann, R.
- CS German Cancer Research Center (DKFZ), Heidelberg, Germany
- SO Journal of Peptide Research (2002), 59(3), 105-114 CODEN: JPERFA; ISSN: 1397-002X
- PB Blackwell Munksgaard
- DT Journal
- LA English
- AΒ The increasing demand for short- and medium-sized peptides in many fields of biol., medical and pharmaceutical research requires optimized and universally applicable high-throughput synthesis and purification techniques at the low-µmol scale. Here, we describe a continuous peptide synthesis/purification approach using the 96-well format. First, a µmol scale peptide synthesis on resin beads was optimized on a novel miniaturized 96-reaction vessel block employing standard Fmoc/tBu-chemical Almost 90% of the synthesized peptides contained the target sequence as the main component, as judged from matrix-assisted laser desorption/ ionization (MALDI) mass spectra. Impurities were mostly related to partially protected peptides. Second, we tested the applicability of ion pair reversed-phase solid-phase extraction (IP-RP-SPE) to purify individual peptides. Depending on the length and predicted hydrophobicity of the peptides, elution was performed with 25 or 35% aqueous acetonitrile in the presence of 0.1% trifluoroacetic acid (TFA). scavengers used during TFA cleavage and partially protected peptides carrying very hydrophobic protecting groups were effectively removed. Using a narrow step gradient, the target peptides were even separated from deleted sequences and protected peptides with similar hydrophobicities. Third, we combined the µmol-scale synthesis in the 96-well format with purification by IP-RP-SPE on a 96-well micro-extraction

plate
format. This simple, fast and parallel approach was tested on 12-mer and
15-mer peptides to map epitopes of T- and B-cell clones, resp. Approx.
80% of all peptides were obtained at purities >90% without purification by
RP-HPLC. In summary, this novel approach has several advantages: (i) the
µmol-scale reduced the cost of peptide synthesis, (ii) large nos. of
peptides were purified faster, (iii) the vols. of eluents and waste were
significantly reduced, and (iv) the RP-HPLC column was not contaminated
with hydrophobic impurities.

RE.CNT 33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L4 ANSWER 9 OF 17 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 2001:417818 CAPLUS
- DN 135:195741
- Site-specific insertion of the (5R*) and (5S*) diastereoisomers of $1-[2-deoxy-\beta-D-erythro-pentofuranosyl]-5-hydroxyhydantoin into oligodeoxyribonucleotides$
- AU Muller, Evelyne; Gasparutto, Didier; Lebrun, Colette; Cadet, Jean
- CS Laboratoire des Lesions des Acides Nucleiques, Service de Chimie

Inorganique et Biologique, UMR 5046, Departement de Recherche Fondamentale sur la Matiere Condensee, CEA-Grenoble, Grenoble, 38054, Fr. SO European Journal of Organic Chemistry (2001), (11), 2091-2099 CODEN: EJOCFK; ISSN: 1434-193X PΒ Wiley-VCH Verlag GmbH DTJournal English LACASREACT 135:195741 OS The insertion of the (5R*) and (5S*) diastereoisomers of AB $1-[2-deoxy-\beta-D-erythro-pentofuranosyl]-5-hydroxyhydantoin- a major$ oxidation product of 2'-deoxycytidine upon exposure to OH radicals, excited photosensitizers, or ozone - into oligonucleotides is reported. It was achieved by means of phosphoramidite chemical, using the solidphase synthesis approach. The synthesis of the phosphoramidite synthon required 6 steps from 3'-O-(tertbutyldimethylsilyl)-2'-deoxycytidine and involved protection of the secondary hydroxy group (5-OH) of the modified base by the nonstandard levulinyl group. This modified phosphoramidite synthon was efficiently incorporated into several oligonucleotides (3-mer, 14-mer, 22-mer) by solid-support assembling. The presence and the integrity of the (5R*) and (5S*) diastereoisomers of 1-[2-deoxy-β-D-erythro-pentofuranosyl]-5hydroxyhydantoin in the synthetic oligomers was confirmed by electrospray ionization mass spectrometry, together with HPLC and MALDI -TOF mass-spectrometric analyses of enzymic digestions. The use of exonucleases (calf spleen phosphodiesterase and bovine intestinal mucosa phosphodiesterase) clearly showed that enzymic hydrolysis of the phosphodiester bonds between the (5R*) and (5S*) diastereoisomers of 1-[2-deoxy-β-D-erythro-pentofuranosyl]-5-hydroxyhydantoin and normal 2'-deoxyribonucleosides is prevented, while endonuclease (nuclease P1) is able to cleave the lesion residue from the oligonucleotides. THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 26 ALL CITATIONS AVAILABLE IN THE RE FORMAT L4ANSWER 10 OF 17 MEDLINE on STN AN2002071557 MEDLINE PubMed ID: 11798020 DN Solid phase synthesis of hydrophobic ΤI difficult sequence peptides on BDDMA-PS support. ΑU Ajikumar P K; Devaky K S School of Chemical Sciences, Mahatma Gandhi University, Kottayam, Kerala, CS India. Journal of peptide science : an official publication of the European SO Peptide Society, (2001 Dec) 7 (12) 641-9. Journal code: 9506309. ISSN: 1075-2617. CY England: United Kingdom DΤ Journal; Article; (JOURNAL ARTICLE) LA English FS Priority Journals EM200206 ED Entered STN: 20020125 Last Updated on STN: 20020627 Entered Medline: 20020626 This article illustrates the successful and efficient solid phase assembly AB of hydrophobic difficult sequence peptides following both t-Boc and Fmoc chemistry. The peptides were synthesized on an optimized 1,4-butanediol dimethacrylate-crosslinked polystyrene support (BDDMA-PS). Four difficult sequence test peptides, VAVAG, VIVIG, QVGQVELG and VQAAIDYING, were synthesized in relatively good yield and purity without any aggregation problems. The peptides were assembled on chloromethylated and 4-hydroxymethylphenoxymethyl (HMP) BDDMA-PS resins. The peptides were fabricated using Boc amino acid 1-hydroxybenzotriazolyl and Fmoc amino acid pentafluorophenyl active esters in coupling reactions. The peptides after synthesis were cleaved from the polymeric support by

exposing the peptidyl resin to 90% trifluroacetic acid/5% thioanisole/5%

EDT mixture. The HPLC and MALDI TOF MS studies of the peptides revealed the high homogeneity of the synthesized peptides. Chloromethylated resin having a functional group loading of 1.14 mmol Cl/g was used for the synthesis. The yield and homogeneity of these peptides synthesized using the new support were high when compared with the conventional DVB-PS resin.

- L4 ANSWER 11 OF 17 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 2003:612955 CAPLUS
- DN 140:321573
- TI Novel methodology for polymer supported oligosaccharide synthesis
- AU Manabe, Shino; Ando, Hiromune; Hanashima, Shinya; Nakahara, Yoshiaki; Ito, Yukishige
- CS RIKEN, The Institute of Physical and Chemical Research, Tokai University, Japan
- SO Tennen Yuki Kagobutsu Toronkai Koen Yoshishu (2001), 43rd, 43-48 CODEN: TYKYDS
- PB Nippon Kagakkai
- DT Journal; General Review
- LA Japanese
 - A review. Solid phase synthesis is now widely recognized as the edge technol. for rapid and efficient oligosaccharide construction. However, it has several disadvantages which should be overcome, (i) the reduced reactivity of substrates, (ii) the difficulty of teal-time reaction monitoring, (iii) limitations on ability of scale-up reactions, and (iv) purification of the desired compds. We choose the PEG (Ave M. W. 550) as a polymer support. It gives homogeneous conditions in reaction mixture, so the reactivity of substrate bound to PEG does not diminished. The scale up of the reaction is possible because the low mol. weight of PEG. Due to its high polarity of PEG, purification of PEG bound sugar was quite simple using silica gel column chromatog. Using the nitro group introduced linker, which is quite stable under typical glycosylation reactions, the oligosaccharide was synthesized on PEG. monitoring of the glycosylation reaction was performed by MALDI -TOF MAS based on the characteristic signal pattern which derives from normal distribution of PEG chain length. The reaction monitoring of deprotection of chloroacetyl group was performed by colorimetric assay by use of (p-nitrobenzyl)pyridine. Chloroacetyl group was selectively reacts with (p-nitrobenzyl)pyridine to give red color under basic conditions. The reaction was semi-quantified by use of NIH Image software. "Catch and release strategy" for the purification of polymer supported oligosaccharide was developed. Solid phase bound cysteine captures the glycosylated product having the chloroacetyl group. Cleavage reaction of Fmoc group releases the sugar via intramol. cyclization process into the solution phase. By repetition of glycosylation/capture/release cycle, the poly(lactosamine) was synthesized on polymer support.
- L4 ANSWER 12 OF 17 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 2000:845660 CAPLUS
- DN 134:116227
- TI Preparation of protected peptides by gel-phase synthesis on butanediol dimethacrylate cross-linked polystyrene support
- AU Roice, M.; Pillai, V. N. Rajasekharan
- CS School of Chemical Sciences, Mahatma Gandhi University, Kottayam, 686 560, India
- SO Protein and Peptide Letters (2000), 7(6), 365-372 CODEN: PPELEN; ISSN: 0929-8665
- PB Bentham Science Publishers
- DT Journal
- LA English
- AB Preparation of fully protected peptide C-terminal esters in high yield and purity by making use of gel-phase synthesis on chloromethyl butanediol dimethacrylate (BDODMA) crosslinked polystyrene is described. The C-terminal amino acid of the peptide was incorporated by cesium salt

method and the step-wise synthesis was carried out using HOBt active ester coupling procedure. The protected peptides were **cleaved** from the support by trans-esterification. The crude peptides were purified by HPLC and characterized by amino acid anal., tlc and **MALDI** TOF MS.

RE.CNT 20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 13 OF 17 MEDLINE on STN

DUPLICATE 1

AN 2000143276 MEDLINE

DN PubMed ID: 10681041

- TI Purification of the c-erbB2/neu membrane-spanning segment: a hydrophobic challenge.
- AU Goetz M; Rusconi F; Belghazi M; Schmitter J M; Dufourc E J
- CS Institut Europeen de Chimie et de Biologie, Ecole Polytechnique, Universites de Bordeaux I et II, B.P. 108, Talence, France.. michael.goetz@iecb-polytechnique.u-bordeaux.fr
- Journal of chromatography. B, Biomedical sciences and applications, (2000 Jan 14) 737 (1-2) 55-61.

 Journal code: 9714109. ISSN: 1387-2273.

CY Netherlands

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 200003

- ED Entered STN: 20000314 Last Updated on STN: 20000314 Entered Medline: 20000301
- AΒ High quality purification of membrane-spanning peptides and proteins remains a challenging problem. In this work we describe a tailored chromatographic purification of a synthetic 35-residue peptide corresponding to the transmembrane region of the tyrosine kinase receptor c-erb2/neu. Composed to over 70% by the amino acids alanine, isoleucine, leucine, phenylalanine and valine, this peptide presents a very hydrophobic character. Product isolation from the complex peptide mixture, obtained after acid cleavage of the resin matrix used during the solid-phase synthesis, represents a difficult task. We propose a three step strategy based on gel permeation and reversed-phase high-performance liquid chromatography, using aprotic polar solvent mixtures. The challenge consisted in obtaining a sufficient amount of an extremely pure sample, in order to allow structural analysis by NMR spectroscopy. Keeping trace of the synthetic peptide throughout the different purification steps was assured by MALDI TOF mass spectrometry, and the final product purity was checked by coupled liquid chromatography-ESI TOF mass spectrometry.
- L4 ANSWER 14 OF 17 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN DUPLICATE 2
- AN 1997:511965 BIOSIS
- DN PREV199799811168
- TI Monitoring the **solid phase synthesis** of analogues of lysobactin and the katanosins using in situ **MALDI** -TOF MS.
- AU Egner, Bryan J.; Bradley, Mark [Reprint author]
- CS Dep. Chem., Univ. Southampton, Highfield, Southampton SO17 1BJ, UK
- SO Tetrahedron, (1997) Vol. 53, No. 41, pp. 14021-14030. CODEN: TETRAB. ISSN: 0040-4020.
- DT Article
- LA English
- ED Entered STN: 10 Dec 1997 Last Updated on STN: 27 Jan 1998
- AB A method of solid phase reaction analysis is described using an in situ cleavage process (TFA vapour) followed by MALDI-TOF MS analysis. The process is demonstrated by the solid

phase synthesis of a depsipeptide based on the antibiotic Lysobactin.

- L4 ANSWER 15 OF 17 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1997:528598 CAPLUS
- DN 127:162115

English to English

- TI Methods for the Chemical Synthesis and Readout of Self-Encoded Arrays of Polypeptide Analogs
- AU Dawson, Philip E.; Fitzgerald, Michael C.; Muir, Tom W.; Kent, Stephen B.
- CS Scripps Research Institute, La Jolla, CA, 92037, USA
- SO Journal of the American Chemical Society (1997), 119(34), 7917-7927 CODEN: JACSAT; ISSN: 0002-7863
- PB American Chemical Society
- DT Journal
- LA English
- AB The synthesis of defined arrays of peptide analogs in conjunction with a simple self-encoded chemical readout system provides a powerful method for the systematic investigation of the relationship between peptide mol. structure and function. A novel **solid-phase**

synthesis procedure was used to prepare arrays of peptide analogs in which a specific thioester modification was systematically incorporated into a unique position in a peptide sequence. The synthesis was carried out in such a way that the resulting arrays contained a defined family of modified peptides, with each peptide mol. containing only a single specific modification. The array of peptide analogs was self-encoded in a positional fashion by incorporating a selectively cleavable thioester bond into the analog structure. Following cleavage of the peptide analog array, anal. of the resulting peptide fragments by MALDI mass spectrometry defined, in a single step, the presence and identity of each peptide analog in the mixture. The feasibility of this approach was demonstrated by the synthesis and mass spectrometric readout of an array of 9 analogs of the 58-residue polypeptide chain of the cCrk N-terminal SH3 domain, before and after folding and affinity selection.

- L4 ANSWER 16 OF 17 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN DUPLICATE 3
- AN 1997:447943 BIOSIS
- DN PREV199799747146
- TI Direct monitoring of organic reactions on polymeric supports.
- AU Carrasco, Michael R. [Reprint author]; Fitzgerald, Michael C.; Oda, Yoshiya; Kent, Stephen B. H.
- CS Scripps Res. Inst., 10550 N. Torrey Pines Rd., La Jolla, CA 92037, USA
- SO Tetrahedron Letters, (1997) Vol. 38, No. 36, pp. 6331-6334. CODEN: TELEAY. ISSN: 0040-4039.
- DT Article
- LA English
- ED Entered STN: 27 Oct 1997
 - Last Updated on STN: 27 Oct 1997
- AB A method to use matrix-assisted laser desorption/ionization mass spectrometry (MALDI-MS) for real-time monitoring of organic reactions on polymeric supports used in solid-phase synthesis is described. The strategy utilizes a synthetic construct that allows for the rapid and convenient direct MALDI analysis of the attached substrates as well as their subsequent chemical cleavage as desired. We have used this strategy to monitor nucleophilic substitutions, palladium-catalyzed coupling reactions, and solid phase peptide synthesis reactions using both Boc- and Fmoc-based chemistries.
- L4 ANSWER 17 OF 17 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1997:285866 CAPLUS
- DN 127:5331
- TI Rapid semi-online monitoring of Fmoc solid-phase peptide synthesis by

matrix-assisted laser desorption/ionization mass spectrometry Talbo, Gert; Wade, John D.; Dawson, Nicola; Manoussios, Mary; Tregear, ΑU Geoffrey W.

CS Howard Florey Institute of Experimental Physiology and Medicine, University of Melbourne, Parkville, VIC 3052, Australia

SO Letters in Peptide Science (1997), 4(2), 121-127 CODEN: LPSCEM; ISSN: 0929-5666

PΒ ESCOM

DTJournal

LAEnglish

AB A simple yet highly effective application of matrix-assisted laser desorption/ionization mass spectrometry (MALDI-MS) for the rapid monitoring of 9-fluorenylmethoxycarbonyl (Fmoc) solid-phase peptide synthesis is described. A few beads of the resin are removed at any desired step during synthesis, the fully protected peptide is cleaved from the resin and an MS spectrum of the analytes present is produced. Some standard side-chain protecting groups may be cleaved off during sample preparation for MS anal.; however, these cleavages are readily identified. Using this approach, incomplete amino acid acylations are readily detected in approx. the same time as by traditional tests such as ninhydrin. The semi-online method also lends itself to ready optimization of synthesis protocols and to the examination of resin-bound peptide side reactions which may not be detectable by chemical means.

THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 15 ALL CITATIONS AVAILABLE IN THE RE FORMAT

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                              AU9SH/BI
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148 AUA1/BI
5 AUA2/BI
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1 AUA6/BI
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1 CARR ROCHELLE/AU
6 CARR RODERICK A/AU
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=> s e3 or e4 or 36 or e7

1159817 "CARR ROBIN"/AU OR "CARR ROBIN A E"/AU OR 36 OR "CARR ROBIN E"/AU

=> s e3-e7

51 ("CARR ROBIN"/AU OR "CARR ROBIN A E"/AU OR "CARR ROBIN ARTHUR"/A U OR "CARR ROBIN ARTHUR ELLIS"/AU OR "CARR ROBIN E"/AU)

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             0 L6 AND MALDI
=> S 16 and spectrometry
             3 L6 AND SPECTROMETRY
≈> dup rem 16
PROCESSING COMPLETED FOR L6
             39 DUP REM L6 (12 DUPLICATES REMOVED)
=> S 19 and spectrometry
             2 L9 AND SPECTROMETRY
=> d bib abs 110 1-2
L10 ANSWER 1 OF 2 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN
     1999:207986 BIOSIS
ΑN
DN
     PREV199900207986
TI
     A photolabile carbamate based dual linker analytical construct for facile
     monitoring of solid phase chemistry: 'TLC' for solid phase?.
TIA
     McKeown, Stephen C. [Reprint author]; Watson, Stephen P.; Carr, Robin
     A. E.; Marshall, Peter
CS
     Discovery Chemistry Unit, Glaxo Wellcome Medicines Research Centre,
     Gunnels Wood Road, Stevenage, Hertfordshire, SG1 2NY, UK
     Tetrahedron Letters, (March 19, 1999) Vol. 40, No. 12, pp. 2407-2410.
SO
     print.
     CODEN: TELEAY. ISSN: 0040-4039.
DT
     Article
     English
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m LA}
ED
     Entered STN: 26 May 1999
     Last Updated on STN: 26 May 1999
AB
     A dual linker analytical construct based on a photolabile carbamate is
     described. Photochemical cleavage from the solid support can be effected
     to afford an analytical fragment, containing the substrate, which is
     sensitised to electrospray mass spectrometry. We believe this
     simple construct now renders all substrates visible to high throughput
     mass specroscopic analysis.
L10 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2004 ACS on STN
ΔN
     2001:64373 CAPLUS
DN
     134:280389
TI
     Single bead characterization using analytical constructs: Application to
     quality control of libraries
UA
     Lorthioir, Olivier; Carr, Robin A. E.; Congreve, Miles S.;
     Geysen, Mario H.; Kay, Corinne; Marshall, Peter; McKeown, Stephen C.;
     Parr, Nigel J.; Scicinski, Jan J.; Watson, Stephen P.
     Glaxo SmithKline Medicines Research Centre, Stevenage Hertfordshire, SG1
CS
     2NY, UK
     Analytical Chemistry (2001), 73(5), 963-970 CODEN: ANCHAM; ISSN: 0003-2700
SO
PΒ
     American Chemical Society
DT
     Journal
LA
     English
AΒ
     Anal. construct technol. was successfully applied to the single-bead anal.
     of a split-mix combinatorial library. Substrates can be released from the
     resin by conventional cleavage for biol. screening. Alternatively, for
     the purpose of anal. and quality control, cleavage at an orthogonal
     construct linker produces an anal. fragment incorporating the substrate.
     This anal. fragment is highly sensitized to electrospray mass
     spectrometry (ESI-MS) and is easily identified by isotope
     labeling. The construct cleavage rendered readily visible even those
     compds. that clearly could not be seen by conventional cleavage and mass
     spectrometry anal. A 1H NMR control experiment proved that the compds.
     cleaved conventionally were, however, present in the sample in good yield
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1-1-1-1

and purity. In view of the data obtained, the authors think that this is a significant and important step toward solving the authors' current quality control problems.

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RE.CNT 26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L1 231 S MALDI AND (SOLID(W) PHASE(W) SYNTHESIS)

L2 7987 S (CLEAVABLE OR CLEAVED OR CLEAVE OR CLEAVAGE) AND (IONIZATION

L3 22 S L1 AND L2

L4 17 DUP REM L3 (5 DUPLICATES REMOVED) E AU=CARR, ROBIN

E CARR ROBIN/AU

L5 1159817 S E3 OR E4 OR 36 OR E7

L6 51 S E3-E7

L7 0 S L6 AND MALDI

L8 3 S L6 AND SPECTROMETRY

L9 39 DUP REM L6 (12 DUPLICATES REMOVED)

L10 2 S L9 AND SPECTROMETRY

FILE 'STNGUIDE' ENTERED AT 21:32:53 ON 11 JUL 2004

FILE 'MEDLINE, BIOSIS, EMBASE, SCISEARCH, CAPLUS, WPIDS' ENTERED AT 21:35:32 ON 11 JUL 2004

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e for e

- L9 ANSWER 1 OF 39 MEDLINE on STN
- TI A 'rule of three' for fragment-based lead discovery?.
- L9 ANSWER 2 OF 39 MEDLINE on STN DUPLICATE 1
- TI Oxidation state of the active-site cysteine in protein tyrosine phosphatase 1B.
- L9 ANSWER 3 OF 39 MEDLINE on STN DUPLICATE 2
- TI Structure-based screening of low-affinity compounds.
- L9 ANSWER 4 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI High-throughput X-ray crystallography and fragment-based screening for drug discovery
- L9 ANSWER 5 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of chemical constructs containing anthracenyl or dansyl groups as UV chromophores
- L9 ANSWER 6 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Single bead characterization using analytical constructs: Application to quality control of libraries
- L9 ANSWER 7 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Reporter Resins for Solid-Phase Chemistry
- L9 ANSWER 8 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Solid-Phase Development of a 1-Hydroxybenzotriazole Linker for Heterocycle Synthesis Using Analytical Constructs
- L9 ANSWER 9 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Amine releasing dual linker analytical constructs for facile monitoring of solid phase chemistry
- L9 ANSWER 10 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of chemical constructs for monitoring reactions on solid supports
- L9 ANSWER 11 OF 39 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN
- TI Correction of Previews 200000521773. Synthesis and SAR of new 5-phenyl-3-ureido-1,5-benzodiazepines as cholecystokinin-B receptor antagonists. Correction of author name.).
- L9 ANSWER 12 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Synthesis and SAR of New 5-Phenyl-3-ureido-1,5-benzodiazepines as Cholecystokinin-B Receptor Antagonists. [Erratum for 2000, Volume 43]
- L9 ANSWER 13 OF 39 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN DUPLICATE 3
- TI Synthesis and SAR of new 5-phenyl-3-ureido-1,5-benzodiazepines as cholecystokinin-B receptor antagonists.

L9 ANSWER 14 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN

1 () **3**

- TI Analysis of solid-phase reactions: product identification and quantification by use of UV-chromophore-containing dual-linker analytical constructs
- L9 ANSWER 15 OF 39 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN DUPLICATE 4
- TI Rapid reaction scanning of solid phase chemistry using resins incorporating analytical constructs.
- L9 ANSWER 16 OF 39 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN DUPLICATE 5
- TI A photolabile carbamate based dual linker analytical construct for facile monitoring of solid phase chemistry: 'TLC' for solid phase?.
- L9 ANSWER 17 OF 39 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN DUPLICATE 6
- TI Cyclic-fused azomethine-, imidate-, and thioimidate methylides: An efficient regiocontrolled entry into spiro-fused pyrrolidines.
- L9 ANSWER 18 OF 39 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN DUPLICATE 7
- TI A short dipolar cycloaddition approach to gamma-lactam alkaloids from Cynometra hankei.
- L9 ANSWER 19 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Hetero-1,3-dipolar cycloadditions of dithiolane-isocyanate imminium methylides: a novel route to 1,3-oxazolidine- and thiazolidine-2-thiones
- L9 ANSWER 20 OF 39 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN DUPLICATE 8
- TI Discovery of 1,5-benzodiazepines with peripheral cholecystokinin (CCK-A) receptor agonist activity: 1. Optimization of the agonist "trigger".
- L9 ANSWER 21 OF 39 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN DUPLICATE 9
- TI Dithiolane-isocyanate iminium methylides: A rapid stereoselective entry into gamma-lactams.
- L9 ANSWER 22 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Enolate bromination in 2-acyl-1,3-dithiane 1-oxides
- L9 ANSWER 23 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of 1,5-benzodiazepine derivatives as cholecystokinin and/or gastrin antagonists
- L9 ANSWER 24 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of 1,5-benzodiazepine-2,4-dione derivatives as cholecystokinin A receptor agonists.
- L9 ANSWER 25 OF 39 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN DUPLICATE 10
- TI Enantioselective synthesis of (R) (-) 2, 6-dimethyl heptanoic acid: The first application of the DITOX asymmetric building block.
- L9 ANSWER 26 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Diastereoselective electrophilic amination of ketone enolates in 2-substituted 2-acyl-1,3-dithiane 1-oxides
- L9 ANSWER 27 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI (Phenylureido)benzodiazepinone antagonists of gastrin and/or cholecystokinin

- L9 ANSWER 28 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI High diastereofacial selectivity in asymmetric Mannich reaction of acyldithiane oxide enolates
- L9 ANSWER 29 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of substituted pyridine insecticidal compounds
- L9 ANSWER 30 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of sulfamylpyridyl alkylsulfonates and analogs as insecticides
- L9 ANSWER 31 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of insecticidal ethers

j () i i

- L9 ANSWER 32 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Fluorinated phenylalkyl ethers, their insecticidal and acaricidal compositions and use, and processes and intermediates for their preparation
- L9 ANSWER 33 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Insecticidal phenylalkyl thioethers, sulfoxides, and sulfones, and processes of their preparation
- L9 ANSWER 34 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of phenylalkenes as intermediates for insecticidal benzyloxypropane derivatives
- L9 ANSWER 35 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Insecticidal (fluoroalkyl)benzene derivatives and processes for their preparation
- L9 ANSWER 36 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Insecticidal and acaricidal arylfluoroalky arylmethyl ethers and processes for their preparation
- L9 ANSWER 37 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Total synthesis of (+)-milbemycin β 3
- L9 ANSWER 38 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Synthesis and reactions of derivatives of 1,7-dioxaspiro[5.5]undec-2-en-4-one
- L9 ANSWER 39 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Total synthesis of (+)-milbemycin β3
- => d bib abs 19 1-10, 14-16
- L9 ANSWER 1 OF 39 MEDLINE on STN
- AN 2003476441 MEDLINE
- DN PubMed ID: 14554012
- TI A 'rule of three' for fragment-based lead discovery?.
- CM Comment on: Drug Discov Today. 2003 Jan 1;8(1):12-6. PubMed ID: 12546981
- AU Congreve Miles; Carr Robin; Murray Chris; Jhoti Harren
- CS Astex Technology Ltd, 436 Cambridge Science Park, Milton Road, CB4 0QA, Cambridge, UK.
- SO Drug discovery today, (2003 Oct 1) 8 (19) 876-7.
 - Journal code: 9604391. ISSN: 1359-6446.
- CY England: United Kingdom
- DT Commentary
 - Journal; Article; (JOURNAL ARTICLE)
- LA English
- FS Priority Journals
- EM 200401
- ED Entered STN: 20031014

Last Updated on STN: 20040129 Entered Medline: 20040128

- L9 ANSWER 2 OF 39 MEDLINE on STN DUPLICATE 1
- AN 2003275676 MEDLINE
- DN PubMed ID: 12802339
- TI Oxidation state of the active-site cysteine in protein tyrosine phosphatase 1B.
- AU van Montfort Rob L M; Congreve Miles; Tisi Dominic; Carr Robin; Jhoti Harren
- CS Astex Technology Ltd, 436 Cambridge Science Park, Milton Road, Cambridge CB4 0QA, UK.
- SO Nature, (2003 Jun 12) 423 (6941) 773-7. Journal code: 0410462. ISSN: 0028-0836.
- CY England: United Kingdom
- DT Journal; Article; (JOURNAL ARTICLE)
- LA English

.

- FS Priority Journals
- OS PDB-10ES; PDB-10ET; PDB-10EU; PDB-10EV
- EM 200307
- ED Entered STN: 20030613 Last Updated on STN: 20030715 Entered Medline: 20030714
- AB Protein tyrosine phosphatases regulate signal transduction pathways involving tyrosine phosphorylation and have been implicated in the development of cancer, diabetes, rheumatoid arthritis and hypertension. Increasing evidence suggests that the cellular redox state is involved in regulating tyrosine phosphatase activity through the reversible oxidization of the catalytic cysteine to sulphenic acid (Cys-SOH). how further oxidation to the irreversible sulphinic (Cys-SO2H) and sulphonic (Cys-SO3H) forms is prevented remains unclear. Here we report the crystal structures of the regulatory sulphenic and irreversible sulphinic and sulphonic acids of protein tyrosine phosphatase 1B (PTP1B), an important enzyme in the negative regulation of the insulin receptor and a therapeutic target in type II diabetes and obesity. We also identify a sulphenyl-amide species that is formed through oxidation of its catalytic cysteine. Formation of the sulphenyl-amide causes large changes in the PTP1B active site, which are reversible by reduction with the cellular reducing agent glutathione. The sulphenyl-amide is a protective intermediate in the oxidative inhibition of PTP1B. In addition, it may facilitate reactivation of PTP1B by biological thiols and signal a unique state of the protein.
- L9 ANSWER 3 OF 39 MEDLINE on STN
- DUPLICATE 2

- AN 2002733649 MEDLINE
- DN PubMed ID: 11983569
- TI Structure-based screening of low-affinity compounds.
- AU Carr Robin; Jhoti Harren
- CS Astex Technology, 250 Cambridge Science Park, Cambridge, UK CB4 OWE.
- SO Drug discovery today, (2002 May 1) 7 (9) 522-7. Ref: 36 Journal code: 9604391. ISSN: 1359-6446.
- CY England: United Kingdom
- DT Journal; Article; (JOURNAL ARTICLE)
 General Review; (REVIEW)
 (REVIEW, TUTORIAL)
- LA English
- FS Priority Journals
- EM 200303
- ED Entered STN: 20021227
 - Last Updated on STN: 20030325 Entered Medline: 20030324
- AB Conventional bioassay-based screening remains a mainstream approach for lead discovery. However, its limitations have meant that other, more biophysical methods, such as X-ray crystallography and NMR, are now being

developed as lead discovery tools. These methods are particularly effective at detecting the binding of low affinity, low molecular weight compounds and transforming them into novel potent leads using structure-guided chemistry. Here, we describe some of the technologies and approaches that are being developed in structure-based screening using X-ray crystallography, which promise to have a major impact on lead discovery.

```
ANSWER 4 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
L9
AN
      2002:744798 CAPLUS
DN
      138:296892
TI
      High-throughput X-ray crystallography and fragment-based screening for
      drug discovery
ΑU
      Carr, Robin
CS
      Astex Technology Ltd., Cambridge, CB4 OWE, UK
      American Genomic/Proteomic Technology (2002), 2(4), 34-36, 38-39
SO
      CODEN: AGTMC7; ISSN: 1537-0003
PΒ
      International Scientific Communications, Inc.
      Journal; General Review
DT
LΑ
      English
      A review. Conventional structure-based drug discovery routinely involves
AB
      the use of crystal structures of target proteins; optimization of lead
      compds. uses three-dimensional information about the protein's binding
      site. Traditionally, x-ray crystallog. has been regarded as a very
      resource-intensive technique, which has restricted its use primarily to
      the lead optimization phase, where it is used to study a small number of
      high-value compds. However, as a result of major technol. advances in
      both software and hardware, rapid determination of protein and protein-ligand
      structures is now allowing x-ray crystallog. to be used in lead screening.
RE.CNT 31
                 THERE ARE 31 CITED REFERENCES AVAILABLE FOR THIS RECORD
                 ALL CITATIONS AVAILABLE IN THE RE FORMAT
      ANSWER 5 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
L9
AN
      2001:265362 CAPLUS
DN
      134:295629
      Preparation of chemical constructs containing anthracenyl or dansyl groups
TI
      as UV chromophores
      Carr, Robin Arthur Ellis; Gehanne, Sylvie; Paio, Alfredo;
      Williams, Geoffrey Martyn; Zaramella, Alessio
PΑ
      Glaxo Group Limited, UK
SO
      PCT Int. Appl., 52 pp.
      CODEN: PIXXD2
DT
      Patent
LA
      English
FAN.CNT 1
      PATENT NO.
                        KIND DATE
                                                   APPLICATION NO. DATE
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      WO 2001025171 Al 20010412
PΙ
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RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
     EP 1218319
                           A1 20020703
                                                  EP 2000-966100 20001003
          R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL
     JP 2003511656
                         T2 20030325
                                                   JP 2001-528123
                                                                         20001003
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PRAI GB 1999-23577

WO 2000-EP9639 W

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20001003

1 6 1 2

Solid phase synthesis and methods of anal. of the products are given. AΒ More specifically "a chemical construct for use in solid phase synthesis comprising a solid support Q having linked thereto via a connecting group Y a substrate R; the connecting group Y having first and second cleavage sites which are orthogonally and selectively cleavable; the second cleavage site being selectively cleavable to release the substrate; and the first cleavage site being located at a position between the second cleavage site and the solid support and being selectively cleavable to release a fragment Fru comprising the substrate and at least a portion of the connecting group Y, wherein the said portion contains a chromophore Cu which facilitates anal. of the fragment Fru by UV, visible or fluorescence spectroscopy, the chromophore Cu having a principal log Emax value of at least 2.5 and wherein (i) the principal log Emax value is at least 1.5 times greater than the principal log Emax of the substrate R; or (ii) the chromophore Cu has an absorption peak at a wavelength remote from absorptions due to the substrate R; and to methods of anal. of products of solid phase synthesis using the constructs". E.g., anthracenes I (X = H, D; R = 3-dimethylaminophenyl, 2-naphthylmethyl, 3-butenyl, tert-butylmethyl) were prepared by solid phase synthesis and analyzed by by UV.

RE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L9 ANSWER 6 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 2001:64373 CAPLUS
- DN 134:280389
- TI Single bead characterization using analytical constructs: Application to quality control of libraries
- AU Lorthioir, Olivier; Carr, Robin A. E.; Congreve, Miles S.; Geysen, Mario H.; Kay, Corinne; Marshall, Peter; McKeown, Stephen C.; Parr, Nigel J.; Scicinski, Jan J.; Watson, Stephen P.
- CS Glaxo SmithKline Medicines Research Centre, Stevenage Hertfordshire, SG1 2NY, UK
- SO Analytical Chemistry (2001), 73(5), 963-970 CODEN: ANCHAM; ISSN: 0003-2700
- PB American Chemical Society
- DT Journal
- LA English
- AB Anal. construct technol. was successfully applied to the single-bead anal. of a split-mix combinatorial library. Substrates can be released from the resin by conventional cleavage for biol. screening. Alternatively, for the purpose of anal. and quality control, cleavage at an orthogonal

construct linker produces an anal. fragment incorporating the substrate. This anal. fragment is highly sensitized to electrospray mass spectrometry (ESI-MS) and is easily identified by isotope labeling. The construct cleavage rendered readily visible even those compds. that clearly could not be seen by conventional cleavage and mass spectrometry anal. A 1H NMR control experiment proved that the compds. cleaved conventionally were, however, present in the sample in good yield and purity. In view of the data obtained, the authors think that this is a significant and important step toward solving the authors' current quality control problems.

RE.CNT 26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L9 ANSWER 7 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 2001:44892 CAPLUS
- DN 134:251943
- TI Reporter Resins for Solid-Phase Chemistry
- AU Congreve, Miles S.; Ladlow, Mark; Marshall, Peter; Parr, Nigel; Scicinski, Jan J.; Sheppard, Tom; Vickerstaffe, Emma; Carr, Robin A. E.
- CS Glaxo Wellcome Cambridge Chemistry Laboratory University Chemical Laboratories, Cambridge, CB2 1EW, UK
- SO Organic Letters (2001), 3(4), 507-510 CODEN: ORLEF7; ISSN: 1523-7060
- PB American Chemical Society
- DT Journal
- LA English
- OS CASREACT 134:251943
- AB An anal. construct resin, designed to aid the anal. of solid-phase chemical, has been mixed in a small proportion with a conventional resin. The anal. construct ("reporter resin") contains two orthogonal linkers that allow cleavage of either the synthetic intermediates (at linker 2) or their anal. enhanced derivs. (at linker 1). The convenient and rapid monitoring of each step in the syntheses of representative library compds. was possible using small resin aliquots.
- RE.CNT 24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L9 ANSWER 8 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 2001:417478 CAPLUS
- DN 135:166813
- TI Solid-Phase Development of a 1-Hydroxybenzotriazole Linker for Heterocycle Synthesis Using Analytical Constructs
- AU Scicinski, Jan J.; Congreve, Miles S.; Jamieson, Craig; Ley, Steven V.; Newman, Emma S.; Vinader, Victoria M.; Carr, Robin A. E.
- CS Department of Chemistry, GlaxoSmithKline Research and Development University Chemical Laboratories, Cambridge, CB2 1EW, UK
- SO Journal of Combinatorial Chemistry (2001), 3(4), 387-396 CODEN: JCCHFF; ISSN: 1520-4766
- PB American Chemical Society
- DT Journal
- LA English
- OS CASREACT 135:166813
- AB The development of a 1-hydroxybenzotriazole linker for the synthesis of heterocyclic derivs. is described, utilizing anal. construct methodol. to facilitate the anal. of resin samples. A UV-chromophore-containing anal. construct enabled the accurate determination of resin loading and the automated monitoring of key reactions using only small quantities of resin. The syntheses of an array of isoxazole derivs. are reported.
- RE.CNT 42 THERE ARE 42 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L9 ANSWER 9 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 2002:46853 CAPLUS
- DN 137:247222
- TI Amine releasing dual linker analytical constructs for facile monitoring of

solid phase chemistry Congreve, Miles S.; Kay, Corinne; Murray, Peter J.; Scicinski, Jan J.; IIA Ley, Steven V.; McKeown, Stephen C.; Watson, Stephen P.; Carr, Robin A. E. Glaxo Wellcome-Cambridge Chemistry Laboratory, Cambridge, CB2 1EW, UK CS Innovation and Perspectives in Solid Phase Synthesis & Combinatorial Libraries: Peptides, Proteins and Nucleic Acids -- Small Molecule Organic Chemistry Diversity, Collected Papers, International Symposium, 6th, York, United Kingdom, Aug. 31-Sept. 4, 1999 (2001), Meeting Date 1999, 217-220. Editor(s): Epton, Roger. Publisher: Mayflower Scientific Ltd., Kingswinford, UK. CODEN: 69CEGV; ISBN: 0-9515735-3-5 DТ Conference LAEnglish Solid phase amine releasing dual linker anal. constructs have been prepared AΒ for reaction monitoring and anal. of chemical stability. The constructs were evaluated against numerous common synthetic reagents to establish their viability for use in solid phase synthesis. THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 11 ALL CITATIONS AVAILABLE IN THE RE FORMAT ANSWER 10 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN L9 AN2000:241132 CAPLUS DN132:278732 Preparation of chemical constructs for monitoring reactions on solid TICarr, Robin Arthur Ellis; Gehanne, Sylvie; Kay, Corinne; ΙN McKeown, Stephen Carl; Murray, Peter John; Paio, Alfredo; Scicinski, Jan Josef; Watson, Stephen Paul; Williams, Geoffrey Martyn; Zaramella, Alessio Glaxo Group Limited, UK PAPCT Int. Appl., 110 pp. SO CODEN: PIXXD2 DTPatent English LA FAN.CNT 1 PATENT NO. KIND DATE APPLICATION NO. DATE ---------A2 A3 PIWO 2000020357 20000413 WO 1999-GB3286 19991005 WO 2000020357 20001026 AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG AU 9961121 A1 20000426 AU 1999-61121 19991005 EP 1119529 20010801 Α2 EP 1999-947750 19991005 20030917 EP 1119529 В1 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO JP 2002526512 T2 20020820 JP 2000-574478 19991005 AT 250018 E 20031015 AT 1999-947750 19991005 PT 1119529 Т PT 1999-947750 20040227 19991005 ES 2207286 20040516 ES 1999-947750 T3 19991005 PRAI GB 1998-21655 19981005 Α WO 1999-GB3286 W 19991005 OS CASREACT 132:278732 AΒ Title constructs comprise a solid support having linked thereto via a connecting group a substrate such that the connecting group has first and

second cleavage sites which are orthogonally and selectively cleavable, the second cleavage site being selectively cleavable to release the

.

substrate, and the first cleavage site being located at a position between the second cleavage site and the solid support and being selectively cleavable to release a fragment comprising the substrate and at least a portion of the connecting group characterized in that cleavage at the first cleavage site forms or introduces on the chemical fragment at the first cleavage site a moiety comprising a sensitizing group (such as an amino group) which sensitizes the chemical fragment to instrumental, e.g. mass spectroscopic, anal. Thus, RNHCO(CH2) 30ZCHMeOH (R = resin, Z = 2-methoxy-5-nitro-1,4-phenylene) (preparation given) was condensed with carbonyldiimidazole and the product amidated by PhCD2N(CO2CMe3) CH2NH2 to give, after deprotection, RNHCO(CH2) 30ZCHMeO2CNHCH2CH2NHCD2Ph which was amidated by HO2C(CH2) 30Z1NHFmocC6H3 (OMe) 2-2,4 (Z1 = 1,4-phenylene) to give, after deprotection, N-benzoylation, and photolysis, H2NCH2CH2N(CD2Ph)CO(CH2) 30Z1CH(NHBz)C6H3 (OMe) 2-2,4 (Z1 unchanged). A mass spectrum of the latter was given.

- L9 ANSWER 14 OF 39 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 2000:733809 CAPLUS
- DN 134:17723
- TI Analysis of solid-phase reactions: product identification and quantification by use of UV-chromophore-containing dual-linker analytical constructs
- AU Williams, Geoff M.; Carr, Robin A. E.; Congreve, Miles S.; Kay, Corinne; McKeown, Stephen C.; Murray, Peter J.; Scicinski, Jan J.; Watson, Stephen P.
- CS Glaxo Wellcome-Cambridge Chem. Lab., Univ. Chemical Lab., Cambridge, CB2 1EW, UK
- SO Angewandte Chemie, International Edition (2000), 39(18), 3293-3296 CODEN: ACIEF5; ISSN: 1433-7851
- PB Wiley-VCH Verlag GmbH
- DT Journal
- LA English
- AB The authors report an enhancement to the recently reported technique of product identification which uses two chemical orthogonal linkers separated by an
 - anal. component, in which release of an anal. fragment composed of an amine group, and inclusion of an isotope label for mass-spectra splitting qualities, was improved by the incorporation of a UV chromophore, which allowed components present in a product mixture to be identified by MS and also quickly quantified by key UV wavelengths.
- RE.CNT 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L9 ANSWER 15 OF 39 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN DUPLICATE 4
- AN 1999:373384 BIOSIS
- DN PREV199900373384
- TI Rapid reaction scanning of solid phase chemistry using resins incorporating analytical constructs.
- AU Murray, Peter John; Kay, Corinne; Scicinski, Jan J. [Reprint author]; McKeown, Stephen C.; Watson, Stephen P.; Carr, Robin A. E.
- CS Glaxo Wellcome-Cambridge Chemistry Laboratory, University Chemical Laboratories, Lensfield Road, Cambridge, CB2 1EW, UK
- SO Tetrahedron Letters, (July 23, 1999) Vol. 40, No. 30, pp. 5609-5612. print.

 CODEN: TELEAY. ISSN: 0040-4039.
- DT Article
- LA English
- ED Entered STN: 9 Sep 1999 Last Updated on STN: 9 Sep 1999
- AB Two analytical constructs, based on orthogonally cleavable linkers, are reported which facilitate the mass spectral analysis of solid phase chemistry. The chemical compatibility and orthogonality of the linkers were established in a parallel reaction study using constructs prepared

specifically for the purpose.

- L9 ANSWER 16 OF 39 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN DUPLICATE 5
- AN 1999:207986 BIOSIS
- DN PREV199900207986
- TI A photolabile carbamate based dual linker analytical construct for facile monitoring of solid phase chemistry: 'TLC' for solid phase?.
- AU McKeown, Stephen C. [Reprint author]; Watson, Stephen P.; Carr, Robin A. E.; Marshall, Peter
- CS Discovery Chemistry Unit, Glaxo Wellcome Medicines Research Centre, Gunnels Wood Road, Stevenage, Hertfordshire, SG1 2NY, UK
- SO Tetrahedron Letters, (March 19, 1999) Vol. 40, No. 12, pp. 2407-2410. print.

 CODEN: TELEAY. ISSN: 0040-4039.
- DT Article

s rec Springer

- LA English
- ED Entered STN: 26 May 1999 Last Updated on STN: 26 May 1999
- AB A dual linker analytical construct based on a photolabile carbamate is described. Photochemical cleavage from the solid support can be effected to afford an analytical fragment, containing the substrate, which is sensitised to electrospray mass spectrometry. We believe this simple construct now renders all substrates visible to high throughput mass specroscopic analysis.

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COST IN U.S. DOLLARS	SINCE FILE	TOTAL
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FULL ESTIMATED COST	45.08	152.61
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DATE: Sunday, July 11, 2004

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	L8	carr-robin.in.	0
	L7	carr.in.	6065
	L6	L1 and L5	78
	L5	(cleavable or cleaved or cleave or cleavage) same (ionization or ionizable or ionisable or ionisation)	776
 ;	L4	(cleavable or cleaved or cleave or cleavage) same (ionization adj tag)	0
S q	L3	(cleavable or cleaved or cleave or cleavage) same(ionization adj tag)	0
	L2	(cleavable or cleaved or cleave or cleavage) with (ionization adj tag)	0
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Search Results - Record(s) 1 through 50 of 78 returned.

☐ 1. Document ID: US 20040115694 A1

Using default format because multiple data bases are involved.

L6: Entry 1 of 78

File: PGPB

Jun 17, 2004

PGPUB-DOCUMENT-NUMBER: 20040115694

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20040115694 A1

TITLE: Methods and compositions for determining the sequence of nucleic acid

molecules

PUBLICATION-DATE: June 17, 2004

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47 Van Ness, Jeffrey Claremont CA US Tabone, John C. Bothell WA US Howbert, J. Jeffry Bellevue WA US Mulligan, John T. Seattle WA US

US-CL-CURRENT: 435/6

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims KMC Draw De

☐ 2. Document ID: US 20040106129 A1

L6: Entry 2 of 78

File: PGPB

Jun 3, 2004

PGPUB-DOCUMENT-NUMBER: 20040106129

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20040106129 A1

TITLE: Mass spectrometric methods for biomolecular screening

PUBLICATION-DATE: June 3, 2004

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47 Crook, Stanley T. US Carlsbad CA Griffey, Richard US Vista CA Hofstadler, Steven Oceanside CA US

US-CL-CURRENT: <u>435/6</u>; <u>702/20</u>

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Kindo

3. Document ID: US 20040096907 A1

L6: Entry 3 of 78

File: PGPB

May 20, 2004

PGPUB-DOCUMENT-NUMBER: 20040096907

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20040096907 A1

TITLE: Quantification of beta amyloid

PUBLICATION-DATE: May 20, 2004

INVENTOR-INFORMATION:

STATE COUNTRY RULE-47 CITY NAME

Bohrmann, Bernd Freiburg DΕ US Ziefen Doebeli, Heinz СН СН Ducret, Axel Riehen СН Guentert, Andreas Boeckten

US-CL-CURRENT: 435/7.1

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims NMC Draw. De

☐ 4. Document ID: US 20040077090 A1

L6: Entry 4 of 78

File: PGPB

Apr 22, 2004

PGPUB-DOCUMENT-NUMBER: 20040077090

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20040077090 A1

TITLE: Whole cell engineering by mutagenizing a substantial portion of a starting

genome, combining mutations, and optionally repeating

PUBLICATION-DATE: April 22, 2004

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

Rancho Santa Fe Short, Jay M. CA

US-CL-CURRENT: 435/471; 435/252.3, 435/254.2

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims FMC Draw De

5. Document ID: US 20040033525 A1

L6: Entry 5 of 78

File: PGPB

Feb 19, 2004

PGPUB-DOCUMENT-NUMBER: 20040033525

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20040033525 A1

TITLE: Releasable nonvolatile mass-label molecules

PUBLICATION-DATE: February 19, 2004

INVENTOR-INFORMATION:

CITY STATE COUNTRY RULE-47 NAME Monforte, Joseph A. Berkeley CA US Palo Alto Becker, Christopher H. CA US Pollart, Daniel J. Menlo Park CA US Shaler, Thomas A. Menlo Park CA US

US-CL-CURRENT: 435/6

Full	Titl	<u>ت</u> را	Citation	Front	Renderor D	lassification.	Date	Reference	Sequences	Attachments	Claims	KOMBO	Err
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6. Document ID: US 20030232758 A1

L6: Entry 6 of 78

File: PGPB

Dec 18, 2003

Dec 11, 2003

PGPUB-DOCUMENT-NUMBER: 20030232758

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030232758 A1

TITLE: Immunological methods and compositions for the treatment of Alzheimer's

disease

PUBLICATION-DATE: December 18, 2003

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

St. George-Hyslop, Peter H. Toronto CA
McLaurin, JoAnne Toronto CA

US-CL-CURRENT: 514/12; 435/320.1, 435/325, 435/69.1, 530/324, 536/23.1

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Mills Draw De 7. Document ID: US 20030228274 A1

File: PGPB

PGPUB-DOCUMENT-NUMBER: 20030228274

PGPUB-FILING-TYPE: new

L6: Entry 7 of 78

DOCUMENT-IDENTIFIER: US 20030228274 A1

Record List Display Page 4 of 19

TITLE: Polyamide chains of precise length

PUBLICATION-DATE: December 11, 2003

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

Rose, Keith Geneva CF

US-CL-CURRENT: 424/78.37; 528/297

8. Document ID: US 20030219830 A1

L6: Entry 8 of 78 File: PGPB Nov 27, 2003

PGPUB-DOCUMENT-NUMBER: 20030219830

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030219830 A1

TITLE: Methods of evaluating glycomolecules for enhanced activities

PUBLICATION-DATE: November 27, 2003

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

Venkataraman, Ganesh Bedford MA US Shriver, Zachary Boston MA US Sasisekharan, Ram Cambridge MA US

US-CL-CURRENT: 435/7.1; 435/68.1, 530/395, 536/53

Full	Title	≘ Citation Front	Review Classification	Date	Reference	Sequences	Attachments	Claims	Kumc	Draw. D
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File: PGPB

Nov 20, 2003

PGPUB-DOCUMENT-NUMBER: 20030215425

PGPUB-FILING-TYPE: new

L6: Entry 9 of 78

DOCUMENT-IDENTIFIER: US 20030215425 A1

TITLE: Epitope synchronization in antigen presenting cells

PUBLICATION-DATE: November 20, 2003

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

Simard, John J. L. Northridge CA US Diamond, David C. West Hills CA US

Record List Display Page 5 of 19

US-CL-CURRENT: 424/93.7; 424/85.1, 424/85.2, 435/372, 514/44, 536/23.1

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims 1000C Draw Da

☐ 10. Document ID: US 20030175819 A1

L6: Entry 10 of 78

File: PGPB

Sep 18, 2003

PGPUB-DOCUMENT-NUMBER: 20030175819

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030175819 A1

TITLE: Methods for identifying modulators of apoptosis

PUBLICATION-DATE: September 18, 2003

INVENTOR-INFORMATION:

NAME CITY

STATE

COUNTRY

RULE-47

Reed, John C.

Rancho Santa Fe

CA

US

Guo, Bin

San Diego

CA

US

US-CL-CURRENT: 435/7.2; 424/9.2

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Killic Draw Da

☐ 11. Document ID: US 20030138866 A1

L6: Entry 11 of 78

File: PGPB

Jul 24, 2003

PGPUB-DOCUMENT-NUMBER: 20030138866

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030138866 A1

TITLE: Novel Osp-C derived peptide fragments

PUBLICATION-DATE: July 24, 2003

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

Mathiesen, Marianne Jartved Hellerup DK
Theisen, Michael Frederiksberg C DK
Holm, Arne Holte DK
Ostergaard, Soren Copenhagen N DK

US-CL-CURRENT: <u>435/7.32</u>

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Review Draws Dr

Record List Display Page 6 of 19

☐ 12. Document ID: US 20030135031 A1

L6: Entry 12 of 78 File: PGPB Jul 17, 2003

PGPUB-DOCUMENT-NUMBER: 20030135031

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030135031 A1

TITLE: Purification of polypeptides

PUBLICATION-DATE: July 17, 2003

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

Rose, Keith Geneva CH
Villain, Matteo Geneva CH
Vizzanova, Jean Geneva CH

US-CL-CURRENT: <u>530/417</u>

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☐ 13. Document ID: US 20030129589 A1

L6: Entry 13 of 78 File: PGPB Jul 10, 2003

PGPUB-DOCUMENT-NUMBER: 20030129589

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030129589 A1

TITLE: DNA DIAGNOSTICS BASED ON MASS SPECTROMETRY

PUBLICATION-DATE: July 10, 2003

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

KOSTER, HUBERT LA JOLLA CA US
LOUGH, DAVID M. BERWICKSHIRE CA GB
XIANG, GOUBING SAN DIEGO US

US-CL-CURRENT: 435/6; 422/68.1

Full Title Citation Front Review Classific.	ation Date Reference Sequences Att	achments Claims RMAC Draw.De
☐ 14. Document ID: US 2003009	06281 A1	
L6: Entry 14 of 78	File: PGPB	May 22, 2003

PGPUB-DOCUMENT-NUMBER: 20030096281

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030096281 A1

TITLE: Methods of making glycomolecules with enhanced activities and uses thereof

PUBLICATION-DATE: May 22, 2003

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

Venkataraman, Ganesh Bedford MΑ US Sasisekharan, Ram Cambridge MA US Shriver, Zachary Boston MΑ US

US-CL-CURRENT: 435/6; 435/101, 435/91.2, 536/123, 536/23.1, 702/20

Full	Titl∈	Citation Front	Review Class	sification	Date	Reference	Sequences	Attachments	Claims	KiniC	Dramu D
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L6: Entry 15 of 78 File: PGPB Apr 24, 2003

PGPUB-DOCUMENT-NUMBER: 20030077595

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030077595 A1

TITLE: Methods and compositions for enhancing sensitivity in the analysis of

biological-based assays

PUBLICATION-DATE: April 24, 2003

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47 Van Ness, Jeffrey Seattle WA US Tabone, John C. Bothell CA US Howbert, J. Jeffry Bellevue WA US Mulligan, John T. Seattle WA US

US-CL-CURRENT: 435/6; 205/777.5, 435/7.9

Full Title	Citation Front	Review Classification	Date	Reference	Sequences	Attachments	Claims	KOMO	Drawn De
□ 16.	Document ID	: US 2003004486	54 A1						
	16 of 78			File:	PGPB		Mar	6,	2003

PGPUB-DOCUMENT-NUMBER: 20030044864

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030044864 A1

TITLE: Cellular engineering, protein expression profiling, differential labeling of peptides, and novel reagents therefor

PUBLICATION-DATE: March 6, 2003

Record List Display Page 8 of 19

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47 Short, Jay M. Rancho Santa Fe CA US Latterich, Martin San Diego CA US Wei, Jing San Diego CA US Levin, Michael San Diego CA US

US-CL-CURRENT: 435/7.23; 702/19

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Numb Praid De

☐ 17. Document ID: US 20030022225 A1

L6: Entry 17 of 78

File: PGPB

Jan 30, 2003

PGPUB-DOCUMENT-NUMBER: 20030022225

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030022225 A1

TITLE: Releasable nonvolatile mass-label molecules

PUBLICATION-DATE: January 30, 2003

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47 Monforte, Joseph A. Berkeley CA US Becker, Christopher H. Palo Alto CA US Pollart, Daniel J. Menlo Park CA US Shaler, Thomas A. Menlo Park CA US

US-CL-CURRENT: 435/6; 435/7.1, 525/10, 530/324, 536/23.1

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims 6000 Braw. De

☐ 18. Document ID: US 20030017483 A1

L6: Entry 18 of 78

File: PGPB

Jan 23, 2003

PGPUB-DOCUMENT-NUMBER: 20030017483

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030017483 A1

TITLE: Modulation of molecular interaction sites on RNA and other biomolecules

PUBLICATION-DATE: January 23, 2003

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

Ecker, David J. Encinitas CA US Griffey, Richard Vista CA US

Crooke, Stanley T.	Carlsbad	CA	US
Sampath, Ranga	San Diego	CA	US
Swayze, Eric	Carlsbad	CA	US
Mohan, Venkatraman	Carlsbad	CA	US
Hofstadler, Steven	Oceanside	CA	US

US-CL-CURRENT: <u>435/6</u>; <u>702/20</u>, <u>703/11</u>

☐ 19. Document ID: US 20020182601 A1

L6: Entry 19 of 78

File: PGPB

Dec 5, 2002

PGPUB-DOCUMENT-NUMBER: 20020182601

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20020182601 A1

TITLE: Method and reagents for analyzing the nucleotide sequence of nucleic acids

PUBLICATION-DATE: December 5, 2002

INVENTOR-INFORMATION:

NAME	CITY	STATE	COUNTRY	RULE-47
Sampson, Jeffrey R.	Burlingame	CA	US	
Myerson, Joel	Berkeley	CA	US	
Tsalenko, Anna M.	Chicago	IL	US	
Sampas, Nicholas M.	San Jose	CA	US	
Webb, Peter G.	Menlo Park	CA	US	
Yakhini, Zohar H.	Zikhron Ya'Acov		IL	

US-CL-CURRENT: 435/6; 536/23.2, 536/24.3

Full Title	Citation Front F	Review Classification	Date Reference	≥ Sequences	Attachments	Claims	KMMC	Draw De
☐ 20.	Document ID:	US 20020169282	2 A1					
L6: Entry	20 of 78		File:	PGPB		Nov	14.	2002

PGPUB-DOCUMENT-NUMBER: 20020169282

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20020169282 A1

TITLE: Solid phase native chemical ligation of unprotected or N-terminal cysteine protected peptides in aqueous solution

PUBLICATION-DATE: November 14, 2002

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

Canne, Lynne

Pacifica

Los Gatos

CA US

Kent, Stephen B.H.

Simon, Reyna J.

San Francisco

CA CA US US

US-CL-CURRENT: 530/334

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Rust Draw De

☐ 21. Document ID: US 20020165383 A1

L6: Entry 21 of 78

File: PGPB

Nov 7, 2002

PGPUB-DOCUMENT-NUMBER: 20020165383

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20020165383 A1

TITLE: Tenebrio antifreeze proteins

PUBLICATION-DATE: November 7, 2002

INVENTOR-INFORMATION:

NAME CITY

Kingston

STATE COUNTRY

RULE-47

Liou, Yih-Cherng

Graham, Laurie A.

Kingston Sydenham CA CA

Walker, Virginia K. Davies, Peter L.

Kingston

CA CA

US-CL-CURRENT: <u>536/23.5</u>; <u>435/320.1</u>, <u>435/325</u>, <u>435/69.1</u>, <u>530/350</u>

☐ 22. Document ID: US 20020142955 A1

L6: Entry 22 of 78

File: PGPB

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Rout Draw De

Oct 3, 2002

PGPUB-DOCUMENT-NUMBER: 20020142955

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20020142955 A1

TITLE: Enzyme-cleavable prodrug compounds

PUBLICATION-DATE: October 3, 2002

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

Dubois, Vincent Fleurus CA ΒE Fernandez, Anne Marie Brussels CA ΒE Gangwar, Sanjeev Alameda CA US Lewis, Evan Daly City CA US

Lobl, Thomas J. Foster City CA US Nieder, Matthew H. Burlingame CA US Pickford, Lesley B. Menlo Park US Trouet, Andre Herent BE Yarranton, Geoffrey T. Burlingame US

US-CL-CURRENT: 514/12; 514/13, 514/14, 514/15, 514/16, 514/17, 530/324, 530/326, 530/327, 530/328

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims សան նուսա նա

☐ 23. Document ID: US 20020132975 A1

L6: Entry 23 of 78

File: PGPB

Sep 19, 2002

PGPUB-DOCUMENT-NUMBER: 20020132975

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20020132975 A1

TITLE: Solid phase native chemical ligation of unprotected or N-terminal cysteine

protected peptides in aqueous solution

PUBLICATION-DATE: September 19, 2002

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

Canne, Lynne Pacifica CA US
Kent, Stephen B.H. San Francisco CA US
Simon, Reyna J. Los Gatos CA US

US-CL-CURRENT: 530/324

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims KWIC Draw De

☐ 24. Document ID: US 20020119456 A1

L6: Entry 24 of 78 Fi

File: PGPB Aug 29, 2002

PGPUB-DOCUMENT-NUMBER: 20020119456

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20020119456 A1

TITLE: Methods and compositions for determining the sequence of nucleic acid

molecules

PUBLICATION-DATE: August 29, 2002

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

Ness, Jeffrey Van Seattle WA US

Tabone, John C.
Howbert, J. Jeffry

Bothell Bellevue WA WA US US

Mulligan, John T.

Seattle

WA

US

US-CL-CURRENT: <u>435/6</u>; <u>250/282</u>

Full | Title | Citation | Front | Review | Classification | Date | Reference | Sequences | Attachments | Claims | KintC | Draw, De

☐ 25. Document ID: US 20020102572 A1

L6: Entry 25 of 78

File: PGPB

Aug 1, 2002

PGPUB-DOCUMENT-NUMBER: 20020102572

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20020102572 A1

TITLE: Mass spectrometric methods for biomolecular screening

PUBLICATION-DATE: August 1, 2002

INVENTOR-INFORMATION:

NAME

CITY

STATE

RULE-47

Crooke, Stanley T.

Carlsbad

CA

US

COUNTRY

Griffey, Richard

Vista

CA

US

Hofstadler, Steven

Oceanside

CA

US

US-CL-CURRENT: 435/6; 702/20

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims KiMC Draw Do

☐ 26. Document ID: US 20020086443 A1

L6: Entry 26 of 78

File: PGPB

Jul 4, 2002

PGPUB-DOCUMENT-NUMBER: 20020086443

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20020086443 A1

TITLE: Magnetic in situ dilution

PUBLICATION-DATE: July 4, 2002

INVENTOR-INFORMATION:

NAME

CITY

STATE

COUNTRY

RULE-47

Bamdad, Cynthia C.

Newton

MA

US

US-CL-CURRENT: 436/526

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims RodC Draw De

Record List Display Page 13 of 19

☐ 27. Document ID: US 20020068301 A1

L6: Entry 27 of 78

File: PGPB

Jun 6, 2002

Apr 11, 2002

PGPUB-DOCUMENT-NUMBER: 20020068301

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20020068301 A1

TITLE: CYCLIC PEPTIDE LIBRARIES AND METHODS OF USE THEREOF TO IDENTIFY BINDING

MOTIFS

PUBLICATION-DATE: June 6, 2002

INVENTOR-INFORMATION:

NAME	CITY	STATE	COUNTRY	RULE-47
LAI, HUNG-SEN	BOSTON	MA	US	
YAFFE, MICHAEL B.	SOMERVILLE	MA	US	
SONGYANG, ZHOU	BROOKLINE	MA	US	
CARRAWAY, KERMIT L. III	WATERTOWN	MA	US	
CANTLEY, LEWIS C.	CAMBRIDGE	MA	US	

US-CL-CURRENT: 435/7.1; 435/194

File: PGPB

L6: Entry 28 of 78

PGPUB-DOCUMENT-NUMBER: 20020042112

PGPUB-FILING-TYPE: new DOCUMENT-IDENTIFIER: US 20020042112 A1

TITLE: DNA DIAGNOSTICS BASED ON MASS SPECTROMETRY

PUBLICATION-DATE: April 11, 2002

INVENTOR-INFORMATION:

NAME	CITY	STATE	COUNTRY	RULE-47
KOSTER, HUBERT	LA JOLLA	CA	US	
LITTLE, DANIEL P.	BOSTON	MA	US	
BRAUN, ANDREAS	SAN DIEGO	CA	US	
LOUGH, DAVID M.	BERWICKSHIRE	CA	GB	
XIANG, GUOBING	SAN DIEGO		US	
VAN DEN BOOM, DIRK	HAMBURG		DE	
JURINKE, CHRISTIAN	HAMBURG		DE	
RUPPERT, ANDREAS	LINDEN		DE	

US-CL-CURRENT: 435/174; 435/6, 435/91.53, 536/23.1

Record List Display Page 14 of 19

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims KNAC Draw Do

29. Document ID: US 20010038070 A1

L6: Entry 29 of 78

File: PGPB

Nov 8, 2001

PGPUB-DOCUMENT-NUMBER: 20010038070

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20010038070 A1

TITLE: Multiplex sequence variation analysis of DNA samples by mass spectrometry

PUBLICATION-DATE: November 8, 2001

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

Hausch, Felix Stanford CA US Jaschke, Andres Berlin DE

US-CL-CURRENT: 250/288

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims KMMC Draws De 30. Document ID: US 20010018511 A1

File: PGPB

L6: Entry 30 of 78

Aug 30, 2001

PGPUB-DOCUMENT-NUMBER: 20010018511

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20010018511 A1

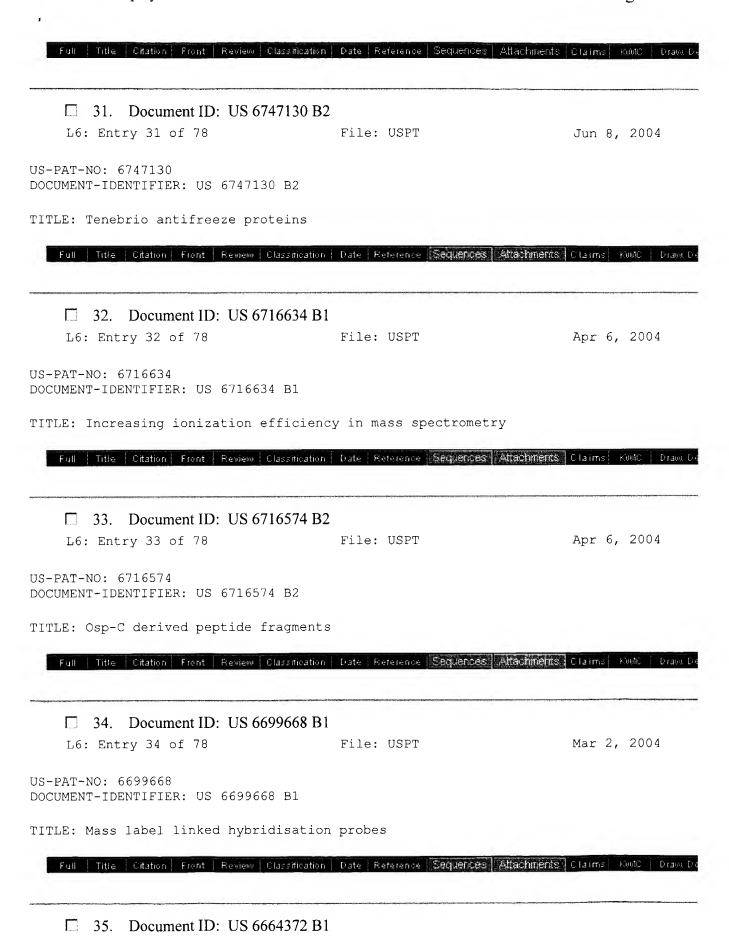
TITLE: INHIBITORS OF THE INTERACTION BETWEEN P53 AND MDM2

PUBLICATION-DATE: August 30, 2001

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47 LANE, DAVID FIFE GB BOTTGER, VOLKER GERMERING DE BOTTGER, ANGELIKA GERMERING DΕ PICKSLEY, STEVEN MICHAEL BRADFORD GB HOCHKEPPEL, HEINZ-KURT **AESCH** СН GARCIA-ECHEVERRIA, CARLOS СН BASEL CHENE, PATRICK MULHOUSE FR FURET, PASCAL THANN FR

US-CL-CURRENT: 536/24.5; 435/375, 435/<u>3</u>77, 435/6, 435/7.1, 530/300, 530/326, 530/327, 530/328, 530/333, 536/24.1



L6: Entry 35 of 78

File: USPT

Dec 16, 2003

US-PAT-NO: 6664372

DOCUMENT-IDENTIFIER: US 6664372 B1

TITLE: Azatide peptidomimetics

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims NOSC Draws Date 36. Document ID: US 6656690 B2

L6: Entry 36 of 78 File: USPT Dec 2, 2003

US-PAT-NO: 6656690

DOCUMENT-IDENTIFIER: US 6656690 B2

TITLE: Mass spectrometric methods for biomolecular screening

Full Title Citation Front Review Classification Data Reference Sequences Attachments Claims ROOC Draws De

37. Document ID: US 6635452 B1

L6: Entry 37 of 78 File: USPT Oct 21, 2003

US-PAT-NO: 6635452

DOCUMENT-IDENTIFIER: US 6635452 B1

** See image for Certificate of Correction **

TITLE: Releasable nonvolatile mass label molecules

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims know Draw Co.

38. Document ID: US 6623928 B2

L6: Entry 38 of 78 File: USPT Sep 23, 2003

US-PAT-NO: 6623928

DOCUMENT-IDENTIFIER: US 6623928 B2

TITLE: Methods and compositions for determining the sequence of nucleic acid

molecules

Full Title Citation Front Review Classification Date Reference **Sequences Stackments** Claims KindC Draw Do

☐ 39. Document ID: US 6613508 B1

L6: Entry 39 of 78

File: USPT

Sep 2, 2003

US-PAT-NO: 6613508

DOCUMENT-IDENTIFIER: US 6613508 B1

TITLE: Methods and compositions for analyzing nucleic acid molecules utilizing

sizing techniques

Full Title Citation Front Review Classification Date Reference **Sequences Attachments** Claims KiMC Draw De

☐ 40. Document ID: US 6602662 B1

L6: Entry 40 of 78

File: USPT

Aug 5, 2003

US-PAT-NO: 6602662

DOCUMENT-IDENTIFIER: US 6602662 B1

TITLE: DNA diagnostics based on mass spectrometry

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Root Draw De

11. Document ID: US 6589485 B2

L6: Entry 41 of 78

File: USPT

Jul 8, 2003

US-PAT-NO: 6589485

DOCUMENT-IDENTIFIER: US 6589485 B2

** See image for Certificate of Correction **

TITLE: Solid support for mass spectrometry

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Killic Draw Do

42. Document ID: US 6558902 B1

L6: Entry 42 of 78 File: USPT May 6, 2003

US-PAT-NO: 6558902

DOCUMENT-IDENTIFIER: US 6558902 B1

TITLE: Infrared matrix-assisted laser desorption/ionization mass spectrometric

analysis of macromolecules

Full Title Citation Front Review Classification Date Reference **Sequences Attachments** Claims Killic Dialog De

☐ 43. Document ID: US 6555692 B1

L6: Entry 43 of 78

File: USPT

Apr 29, 2003

US-PAT-NO: 6555692

DOCUMENT-IDENTIFIER: US 6555692 B1

TITLE: Preparation and use of bifunctional molecules having DNA sequence binding

specificity

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims MMC Draw Dr ☐ 44. Document ID: US 6552167 B1

L6: Entry 44 of 78

File: USPT

Apr 22, 2003

US-PAT-NO: 6552167

DOCUMENT-IDENTIFIER: US 6552167 B1

** See image for Certificate of Correction **

TITLE: Polyamide chains of precise length

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Kimo Draw De ☐ 45. Document ID: US 6506906 B1 L6: Entry 45 of 78 File: USPT Jan 14, 2003

US-PAT-NO: 6506906

DOCUMENT-IDENTIFIER: US 6506906 B1

TITLE: Preparation and use of bifunctional molecules having DNA sequence binding

specificity

Full Title Citation Front Review Classification Date Reference Sequences Affactments Claims Kinto Draw De ☐ 46. Document ID: US 6500621 B2 L6: Entry 46 of 78 File: USPT Dec 31, 2002

US-PAT-NO: 6500621

DOCUMENT-IDENTIFIER: US 6500621 B2

** See image for Certificate of Correction **

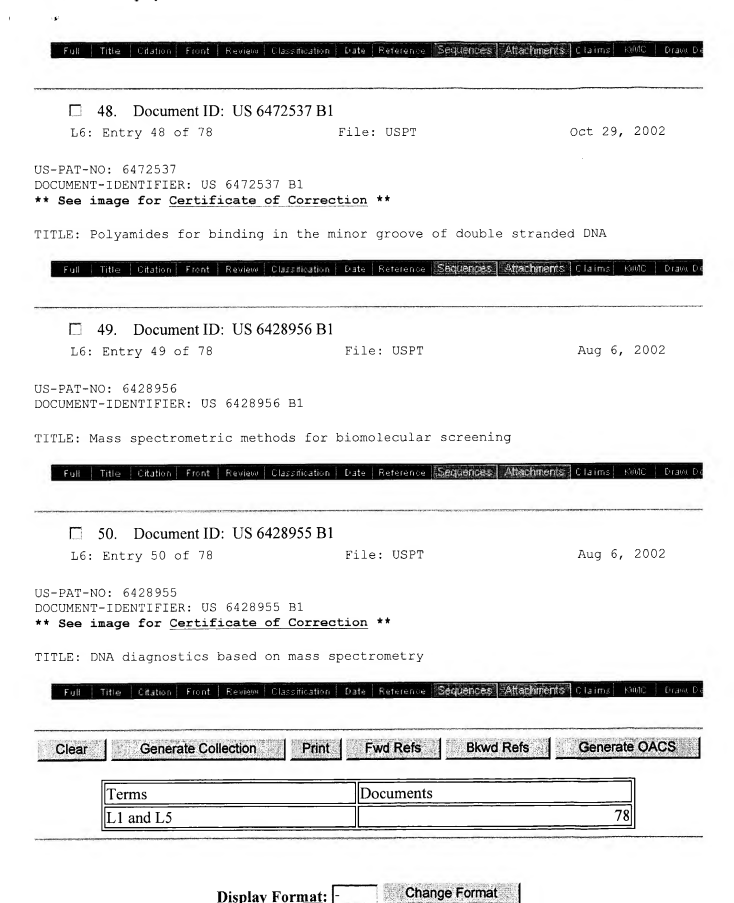
TITLE: DNA diagnostics based on mass spectrometry

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Kinto Drawa Do ☐ 47. Document ID: US 6495314 B1 L6: Entry 47 of 78 File: USPT Dec 17, 2002

US-PAT-NO: 6495314

DOCUMENT-IDENTIFIER: US 6495314 B1

TITLE: Process for characterizing proteins



WEST Search History

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DATE: Sunday, July 11, 2004

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	DB=PC	GPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD; PLUR=YES; OP=OR	
	L6	L1 and L5	78
	L5	(cleavable or cleaved or cleave or cleavage) same (ionization or ionizable or ionisable or ionisation)	776
	L4	(cleavable or cleaved or cleave or cleavage) same (ionization adj tag)	0
	L3	(cleavable or cleaved or cleave or cleavage) same(ionization adj tag)	0
	L2	(cleavable or cleaved or cleave or cleavage) with (ionization adj tag)	0
	L1	MALDI and (solid adj phase adj synthesis)	578

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Search Results - Record(s) 51 through 78 of 78 returned.

51. Document ID: US 6392024 B1

Using default format because multiple data bases are involved.

L6: Entry 51 of 78

File: USPT

May 21, 2002

US-PAT-NO: 6392024

DOCUMENT-IDENTIFIER: US 6392024 B1

TITLE: Tenebrio antifreeze proteins

DATE-ISSUED: May 21, 2002

INVENTOR-INFORMATION:

CITY STATE ZIP CODE COUNTRY NAME Graham; Laurie A. Kingston CA CA Kingston Liou; Yih-Cherng Sydenham CA Walker; Virginia K. Davies; Peter L. Kingston CA

US-CL-CURRENT: 536/23.5; 435/252.3, 435/254.11, 435/254.21, 435/254.22, 435/320.1, 435/6, 536/23.1



52. Document ID: US 6387628 B1

L6: Entry 52 of 78

File: USPT

May 14, 2002

US-PAT-NO: 6387628

DOCUMENT-IDENTIFIER: US 6387628 B1

** See image for <u>Certificate of Correction</u> **

TITLE: Mass spectrometric detection of polypeptides



☐ 53. Document ID: US 6329146 B1

L6: Entry 53 of 78

File: USPT

Dec 11, 2001

US-PAT-NO: 6329146

DOCUMENT-IDENTIFIER: US 6329146 B1

Record List Display Page 2 of 8

** See image for Certificate of Correction **

TITLE: Mass spectrometric methods for biomolecular screening



54. Document ID: US 6326468 B1

L6: Entry 54 of 78

File: USPT

Dec 4, 2001

US-PAT-NO: 6326468

DOCUMENT-IDENTIFIER: US 6326468 B1

TITLE: Solid phase native chemical ligation of unprotected or n-terminal cysteine

protected peptides in aqueous solution

Full Title Citation Front Review Classification Date Reference **Sequences Attachneers** Claims KWC Draw De

☐ 55. Document ID: US 6322970 B1

L6: Entry 55 of 78

File: USPT

Nov 27, 2001

US-PAT-NO: 6322970

DOCUMENT-IDENTIFIER: US 6322970 B1

** See image for Certificate of Correction **

TITLE: Mass spectrometric detection of polypeptides

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Kint Dram Do

56. Document ID: US 6312893 B1

L6: Entry 56 of 78 File: USPT Nov 6, 2001

US-PAT-NO: 6312893

DOCUMENT-IDENTIFIER: US 6312893 B1

** See image for Certificate of Correction **

TITLE: Methods and compositions for determining the sequence of nucleic acid

molecules

☐ 57. Document ID: US 6310180 B1

L6: Entry 57 of 78

File: USPT

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims FWC Draw De

Oct 30, 2001

US-PAT-NO: 6310180

DOCUMENT-IDENTIFIER: US 6310180 B1

Record List Display Page 3 of 8

TITLE: Method for synthesis of proteins

Full Title Citation Front Review Classification Date Reference **Sequences Attachments** Claims RMC Draw De

☐ 58. Document ID: US 6300076 B1

L6: Entry 58 of 78

File: USPT

Oct 9, 2001

US-PAT-NO: 6300076

DOCUMENT-IDENTIFIER: US 6300076 B1

TITLE: DNA diagnostics based on mass spectrometry

Full Title Citation Front Review Classification Date Reference **Sequences Affectineris** Claims RMC Draw De

☐ 59. Document ID: US 6277573 B1

L6: Entry 59 of 78

File: USPT

Aug 21, 2001

US-PAT-NO: 6277573

DOCUMENT-IDENTIFIER: US 6277573 B1

** See image for Certificate of Correction **

TITLE: DNA diagnostics based on mass spectrometry

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims KMC Draw De

☐ 60. Document ID: US 6268144 B1

L6: Entry 60 of 78

File: USPT

Jul 31, 2001

US-PAT-NO: 6268144

DOCUMENT-IDENTIFIER: US 6268144 B1

** See image for Certificate of Correction **

TITLE: DNA diagnostics based on mass spectrometry

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Kill Draw De Common Co

L6: Entry 61 of 78

File: USPT

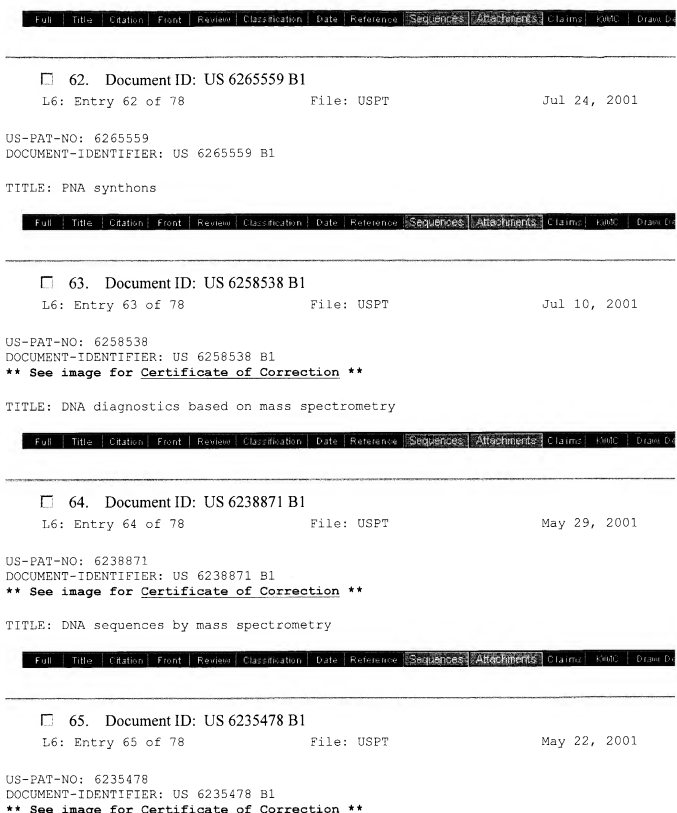
Jul 31, 2001

US-PAT-NO: 6268131

DOCUMENT-IDENTIFIER: US 6268131 B1

** See image for Certificate of Correction **

TITLE: Mass spectrometric methods for sequencing nucleic acids



** See image for Certificate of Correction **

TITLE: DNA diagnostics based on mass spectrometry

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Func Draw De

☐ 66. Document ID: US 6225450 B1

L6: Entry 66 of 78

File: USPT

May 1, 2001

US-PAT-NO: 6225450

DOCUMENT-IDENTIFIER: US 6225450 B1

** See image for Certificate of Correction **

TITLE: DNA sequencing by mass spectrometry

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Root Draw De

☐ 67. Document ID: US 6221605 B1

L6: Entry 67 of 78

File: USPT

Apr 24, 2001

US-PAT-NO: 6221605

DOCUMENT-IDENTIFIER: US 6221605 B1

** See image for Certificate of Correction **

TITLE: DNA diagnostics based on mass spectrometry

Full Title Citation Front Review Classification Date Reference **Sequences Attachments** Claims KMIC Draw De

☐ 68. Document ID: US 6221601 B1

L6: Entry 68 of 78

File: USPT

Apr 24, 2001

US-PAT-NO: 6221601

DOCUMENT-IDENTIFIER: US 6221601 B1

TITLE: DNA diagnostics based on mass spectrometry

Full Title Citation Front Review Classification Date Reference **Sequences Attachments** Claims KiviC Draw Do

☐ 69. Document ID: US 6197498 B1

L6: Entry 69 of 78

File: USPT

Mar 6, 2001

US-PAT-NO: 6197498

DOCUMENT-IDENTIFIER: US 6197498 B1

** See image for Certificate of Correction **

TITLE: DNA diagnostics based on mass spectrometry

Full Title Citation Front Review Classification Date Reference Sequences Attachinents Claims MMC Draw Da

70. Document ID: US 6194144 B1

L6: Entry 70 of 78

File: USPT

Feb 27, 2001

Nov 7, 2000

US-PAT-NO: 6194144

DOCUMENT-IDENTIFIER: US 6194144 B1

** See image for Certificate of Correction **

TITLE: DNA sequencing by mass spectrometry

Full Title Citation Front Review Classification Date Reference **Sequences Attachments** Claims 1996 Draw De 71. Document ID: US 6143932 A

File: USPT

US-PAT-NO: 6143932

DOCUMENT-IDENTIFIER: US 6143932 A

L6: Entry 71 of 78

TITLE: Selectively N-alkylated peptidomimetic combinatorial libraries and compounds

therein

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims KiMC Diam Do ☐ 72. Document ID: US 6063569 A

L6: Entry 72 of 78

File: USPT

May 16, 2000

Mar 28, 2000

US-PAT-NO: 6063569

DOCUMENT-IDENTIFIER: US 6063569 A

TITLE: Methods for automated synthesis of PNA-DNA chimeras and compositions thereof

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims Kimic Diam De 73. Document ID: US 6043031 A

File: USPT

US-PAT-NO: 6043031

DOCUMENT-IDENTIFIER: US 6043031 A

L6: Entry 73 of 78

** See image for Certificate of Correction **

TITLE: DNA diagnostics based on mass spectrometry

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims NAC Draw Da ☐ 74. Document ID: US 6027890 A

L6: Entry 74 of 78

File: USPT

Feb 22, 2000

US-PAT-NO: 6027890

DOCUMENT-IDENTIFIER: US 6027890 A

** See image for Certificate of Correction **

TITLE: Methods and compositions for enhancing sensitivity in the analysis of

biological-based assays

Full Title Citation Front Review Classification Data Reference Sequences Attachments Claims KnotC Draw. De

75. Document ID: US 5830655 A

L6: Entry 75 of 78

File: USPT

Nov 3, 1998

US-PAT-NO: 5830655

DOCUMENT-IDENTIFIER: US 5830655 A

TITLE: Oligonucleotide sizing using cleavable primers

Full Title Citation Front Review Classification Date Reference **Sequences Attachments** Claims KiMC Draw De

☐ 76. Document ID: US 5700642 A

L6: Entry 76 of 78

File: USPT

Dec 23, 1997

US-PAT-NO: 5700642

DOCUMENT-IDENTIFIER: US 5700642 A

TITLE: Oligonucleotide sizing using immobilized cleavable primers

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims KMC Draw, De

77. Document ID: US 5691141 A

L6: Entry 77 of 78

File: USPT

Nov 25, 1997

US-PAT-NO: 5691141

DOCUMENT-IDENTIFIER: US 5691141 A

** See image for Certificate of Correction **

TITLE: DNA sequencing by mass spectrometry

Full Title Citation Front Review Classification Date Reference **Sequences Attachments** Claims MMC Draw De

78. Document ID: US 5547835 A

L6: Entry 78 of 78

File: USPT

Aug 20, 1996

US-PAT-NO: 5547835

DOCUMENT-IDENTIFIER: US 5547835 A

TITLE: DNA sequencing by mass spectrometry

Full	Title Citation Fron	t Review Cla	ssification Dat	te Reference	Sequences Atla	Aments Claims	KMMC Draw, D
Clear	Generate C	ollection	Print	Fwd Refs	Bkwd Refs	Gene	rate OACS
	Terms			Documents	<u> </u>		
	L1 and L5						78

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